

Sensitivity of aerosol properties to new particle formation mechanism and to primary emissions in a continental-scale chemical transport model

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[1] Four theoretical formulations of new particle formation (NPF) and one empirical formulation are used to examine the sensitivity of observable aerosol properties to NPF formulation and to properties of emitted particles in a continental-scale model for the United States over a 1-month simulation (July 2004). For each formulation the dominant source of Aitken mode particles is NPF with only a minor contribution from primary emissions, whereas for the accumulation mode both emissions and transfer of particles from the Aitken mode are important. The dominant sink of Aitken mode number is coagulation, whereas the dominant sink of accumulation mode number is wet deposition (including cloud processing), with a minor contribution from coagulation. The aerosol mass concentration, which is primarily in the accumulation mode, is relatively insensitive to NPF formulation despite order-of-magnitude differences in the Aitken mode number concentration among the different parameterizations. The dominant sensitivity of accumulation mode number concentration is to the number of emitted particles (for constant mass emission rate). Comparison of modeled aerosol properties with aircraft measurements shows, as expected, better agreement in aerosol mass concentration than in aerosol number concentration for all NPF formulations considered. These comparisons yield instances of rather accurate simulations in the planetary boundary layer, with poor model performance in the free troposphere attributed mainly to lack of representation of biomass burning and/or to long-range transport of particles from outside the model domain. Agreement between model results and measurements is improved by using smaller grid cells (12 km versus 60 km).

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1. Introduction

[2] Indirect aerosol forcing, the change in reflected short-wave radiation due to enhancement of cloud drop number concentration and cloud reflectivity by anthropogenic aerosols, remains the largest uncertainty in radiative forcing of climate change over the industrial period [Intergovernmental Panel on Climate Change, 2007]. Recent studies have concluded that the main sources of uncertainty in aerosol indirect effects are the uncertainties in cloud parameterization and in aerosol particle number concentration [Menon et al., 2002; Chen and Penner, 2005]. As global models often employ empirical or physically based parameterizations to relate cloud droplet number concentration to aerosol number concentration [Boucher and Lohmann, 1995; Kiehl et al.,

[3] The principal processes controlling aerosol number concentration are emission of primary particles, new particle formation (NPF) from vapor precursors, transport, coagulation, and wet and dry deposition. Any uncertainty in representation of these processes in models contributes to uncertainty in modeled aerosol number concentration. Additionally changes in aerosol properties by atmospheric chemical and physical processes affect the dynamics of aerosol particles and in turn their number concentration and size distribution. Aerosol particles can grow or shrink by coagulation, condensation, evaporation, and surface reactions. Coagulation directly reduces concentration of aerosol number (but not mass) and although condensation, evaporation, and surface reactions do not directly alter number concentration, coupling to vapor concentrations and particle growth makes these processes important to considerations of sizedependent number concentration. Accurate knowledge of

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^{2000;} Ghan et al., 2001; Menon et al., 2002], accurate knowledge of aerosol number concentration and size distribution is important for considerations of the aerosol-cloud interactions in general and of aerosol indirect effects in particular.

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these processes and representation of their effects in climate models is therefore essential to accurate representation of climate change [Ghan and Schwartz, 2007].

- [4] The influence of different processes on overall aerosol number concentration has previously been examined in a term-by-term timescale analysis of the aerosol dynamic equation under typical urban conditions by *Zhang and Wexler* [2002], who confirmed the intuitive notion that particulate emissions and new particle formation are important determinants of the size distribution of number concentration for urban and regional aerosols. This finding suggests particulate sources as key processes for sensitivity analysis in chemical transport models.
- [5] Atmospheric aerosol particles can be classified according to their sources as either primary—those that are injected directly into the atmosphere, and secondary—those that are formed from nucleation of precursor gaseous substances (although both primary and secondary particles can accrete primary and secondary particulate mass by coagulation, condensation, and reaction, somewhat blurring the distinction between primary and secondary particulate matter). Primary aerosol particles commonly provide the dominant contribution to mass concentrations and tend to be considerably larger than secondary aerosol particles, which often dominate number concentrations.
- [6] Key examples of primary aerosol particles are sea salt, crustal material (i.e., dust), and anthropogenically emitted particles. Emissions of natural aerosols such as dust and sea salt are strongly dependent on wind speed and surface features, and may occur over large areas, whereas anthropogenic aerosols often arise from point sources. Inventories of anthropogenic emissions, generally developed in support of ambient air quality standards, are based on mass rather than number, and number-based inventories are not anticipated in the near future; consequently, there is little information on the numbers or sizes of primary (i.e., emitted) particles in currently available emissions databases. The impact of emissions on the aerosol size distribution has been investigated by *Adams and Seinfeld* [2003], who stressed the need for accurate size-resolved emissions.
- [7] Secondary aerosol particles evolve from stable nuclei that spontaneously form from vapor precursors and subsequently grow by condensation of vapors and coagulation with other such particles. Here a distinction is made between nucleation and new particle formation: nucleation refers to the formation of stable particles, here taken as diameter 1 nm, whereas new particle formation refers to the production of those nucleated particles that have grown by further condensation of sulfuric acid, ammonia, water, and/ or organics or by coagulation with comparably sized particles until they have attained a diameter that is observable by current instrumentation, here taken as 3 nm. The rates of nucleation J and new particle formation J_{NPF} are defined as the number of nucleated particles and new particles, respectively, produced per unit time per unit volume. Because some nuclei are scavenged through coagulation with other nuclei or by larger aerosol particles before attaining the size at which they become new particles, the rate of new particle formation is less than that of nucleation by a factor that depends on the relative importance of the growth rate of the nuclei and the rate at which they are lost to coagulation.

- [8] Various mechanisms and formulations for NPF have been proposed. Three mechanisms considered here are binary homogeneous nucleation (BHN) of sulfuric acid and water, ternary homogeneous nucleation (THN) of sulfuric acid, water, and ammonia, and nucleation by ionion recombination (NIIR). Formulations for BHN and THN yield rates of nucleation J that depend on the concentration of sulfuric acid molecules [H₂SO₄], the relative humidity RH, the temperature T, and, in the case of THN, the dry air molal mixing ratio of ammonia [NH₃]. Nucleation is favored by lower temperatures and higher relative humidities and, in the case of THN, by higher ammonia concentrations, as ammonia neutralizes the sulfuric acid and reduces the energy barrier to nucleation, yielding nucleation rates that can greatly exceed those from BHN under otherwise similar conditions. An alternative mechanism for nucleation is provided by NIIR. According to this mechanism, cosmogenic ions, which are ubiquitous at low concentrations in the atmosphere, can in the presence of typically measured sulfuric acid concentrations lead to the generation of large molecular-ion clusters, which upon recombination with molecular ions of opposite charge yield thermodynamically stable neutral clusters that with further growth can form new particles.
- [9] Nucleation parameterizations contain large uncertainties, and debate continues over the mechanisms that dominate, or even operate, under given conditions. As nucleation rates are highly nonlinear in controlling variables (temperature, relative humidity, and concentrations of precursor gases), accurately modeling nucleation occurrence and rates is challenging, as large spatial inhomogeneities of controlling variables within a single to grid cell (typically tens of kilometers on a side) can result in large differences between average nucleation rates over a grid cell and calculated rates based on average values of these variables. Nonetheless the importance of nucleation as a source of new particles warrants investigation of nucleation, new particle formation, and their consequences.
- [10] Several model studies have examined nucleation and new particle formation using different assumptions and parameterizations of nucleation rates. Adams and Seinfeld [2002], using a global model, made the assumption that nucleation occurred, if at all, only if the sulfuric acid concentration after condensation onto existing particles was taken into account exceeded a critical value that depended only on temperature and relative humidity. Spracklen et al. [2006], in an investigation of the contribution of boundary layer nucleation events to number concentration on regional and global scales, assumed that nucleation above the boundary layer occurred by BHN, and that in the boundary layer it occurred at a rate that is directly proportional to the concentration of sulfuric acid; the effect of the constant of proportionality was further investigated in a global model by Spracklen et al. [2008]. Sotiropoulou et al. [2006] examined air quality and CCN concentrations in a regional model with new particle formation occurring by THN. Parameterizations of new particle formation by binary, ternary, and ion-induced nucleation were examined in a global model by Lucas and Akimoto [2006], who concluded that nucleation parameterizations exhibit large uncertainties arising from inconsistencies between formulations for the same mechanism, numerical

fitting errors, and the like. Yu et al. [2008] investigated the importance of ion-mediated nucleation (NIIR) to number concentration in a global model and identified regions where this mechanism would yield large nucleation rates.

[11] The present study examines the effects of uncertainties in parameterizations of nucleation and new particle formation, as well as in numbers and sizes of emitted particles, on the size distribution of aerosol number concentration and on other aerosol properties. Only anthropogenic sources of primary particles (sulfate, nitrate, organic and elemental carbon, and a further, unspecified, category) are considered in this study; dust, sea salt, and biomass burning (which may be anthropogenic) are not included. Because anthropogenic emission inventories are based on mass rather than number, as noted above, models representing aerosol number concentration therefore need to determine number and size distribution of emissions from massbased inventories until other means become available [Zhang and Wexler, 2004; Zhang et al., 2004]. This situation contributes additional uncertainty to aerosol number concentrations beyond that from the uncertainties in the mass emission inventories themselves. The sensitivity of modeled aerosol number concentrations to this uncertainty in emissions is also examined here.

2. Model Description

[12] The model calculations reported here were carried out using Version 4.4 of the Community Multiscale Air Quality (CMAQ) model [Byun and Ching, 1999; Binkowski, 1999; Binkowski and Roselle, 2003], in which the principal processes affecting aerosol particles that are represented are advection and diffusion, gas-phase chemistry and aqueous sulfate chemistry, aerosol microphysics, aerosol thermodynamics, redistribution of aerosols in convective clouds, and wet and dry deposition.

[13] Gas-phase chemistry is treated using the Statewide Air Pollution Research Center of the University of California at Riverside 1999 (SAPRC99) (W. P. L. Carter, Implementation of the SAPRC-99 chemical mechanism into the models-3 framework, report to the U.S. Environmental Protection Agency, 29 Jan. 2000, available at http://www.cert.ucr.edu/ ~carter/absts.htm#s99mod3) formulation representing volatile organic carbon-nitrogen oxide (VOC-NO_x) chemistry plus oxidation of SO₂ by OH. SAPRC99 as implemented in CMAQ treats 72 species having 214 reactions, involving inorganics (NO_x, O₃, H₂O₂, CO, SO₂, etc.), organics (PANs, 5 lumped alkenes, 2 aromatics, 2 olefins, isoprene, terpenes, etc), and radicals (OH, HO₂, organic peroxy radicals, etc.), of which OH is important as an initiator of sulfuric acid production (as discussed in section 2.2). Aerosol-forming species are water, sulfuric acid, and ammonia (which are treated here as the sole sources of nucleation), nitric acid, and anthropogenic and biogenic secondary organic aerosol (SOA), which forms from precursors (importantly alkanes, alkenes, aromatics, and monoterpenes) at rates that are derived from experiment [Pandis et al., 1992].

[14] Aqueous sulfate chemistry is based on the Regional Acid Deposition Model [*Chang et al.*, 1987], which treats absorption of gases into cloud droplets, dissociation, oxidation of S(IV) to S(VI), and wet deposition for 11 gaseous

species including SO₂, OH, O₃, and H₂O₂, and 12 aerosol species including sulfate, ammonium, and nitrate.

[15] Aerosol microphysics is treated by a modal representation [Binkowski and Roselle, 2003] consisting of three modes (Aitken, accumulation, and coarse) with each mode being characterized by three moments with respect to the dry diameter (i.e., not including water mass). The number concentration in each mode is assumed to have a lognormal distribution, and the three moments are directly proportional to the total concentrations of number, surface area, and volume. Within each mode internal mixing is assumed; that is, all particles within a mode are treated as having the same chemical composition. The moments of the modes are affected by some or all of the following processes: NPF, condensation of sulfuric acid or other gases (besides water vapor), coagulation, and wet and dry deposition. Newly formed particles are treated as entering the Aitken mode, the criterion for which is that subsequent to nucleation the particles have grown to a diameter of 3 nm as discussed above. Condensation can occur in all three modes. Coagulation involving coarse mode particles is assumed to be negligible and is not represented. Aitken mode particles in clouds are assumed to be scavenged by accumulation mode particles (resulting in a decrease in both mass and number concentrations in the Aitken mode, and an increase in mass concentration, but not number concentration, in the accumulation mode) at a constant rate over the cloud lifetime, and unactivated accumulation mode particles can also be scavenged by activated cloud drops (resulting in a reduction of number but not mass concentration in the accumulation mode; this loss is classified as wet deposition rather than coagulation). As new particles grow by condensation of gases and coagulation with other newly formed particles and the mean (dry) diameter of the Aitken mode approaches that of the accumulation mode, in some situations a fraction of each moment of the Aitken mode is transferred to the accumulation mode; this is termed intermodal transfer. This algorithm, discussed below, is formulated here as in the work of Binkowski and Roselle [2003] without modification.

[16] Aerosol thermodynamics is treated by ISORROPIA [Nenes et al., 1998], which calculates the composition and phase state of an ammonium-sulfate-nitrate-chloride-sodium-water inorganic mixture in thermodynamic equilibrium with the corresponding gas-phase precursor species, thus explicitly accounting for the change in size of particles with respect to relative humidity.

[17] The modeling domain covers the continental United States at a resolution of 60 km with 21 vertical levels (based on pressure) extending up to a pressure of 10 hPa (which typically occurs near 30 km); thus the total number of grid cells in the domain is $95 \times 60 \times 21$, or approximately 1.2×10^5 . Processes in the northeast Unites States are examined in greater detail using a nested domain with a resolution of 12 km. The simulation time was chosen to coincide with the campaign period of the International Consortium for Atmospheric Research on Transport and Transformation (ICARTT) project, 1 July to 31 August 2004 [Fehsenfeld et al., 2006] to permit comparison with observations.

[18] For the time integration, CMAQ employs operator splitting, in which several processes are performed separately with different time steps, each being dynamically determined on the basis of numerical stability and on the

level of accuracy required. The overall time interval, which includes an integral number of time steps for each process, is 15 min; model output was written at every 3 h. Thus the 1-month model run contains $\sim 3 \times 10^3$ time intervals, and output was written 248 times.

2.1. New Particle Formation

- [19] Sensitivity of aerosol number concentration to new particle formation was examined by considering several parameterizations for the rates of nucleation, J, and of new particle formation, $J_{\rm NPF}$. In addition, one empirical NPF formulation was considered.
- [20] Two binary homogeneous nucleation parameterizations were used: that of Jaecker-Voirol and Mirabel [1989] (hereinafter referred to as BHNJ) and that of Vehkamäki et al. [2002] (hereinafter referred to as BHNV). The nucleation rate according to BHNJ, J_{BHNJ} , was originally presented graphically and designed for simple interpolation in T and RH. For the present study, graphs of log J_{BHNJ} versus log $[H_2SO_4]$ for T = [223, 248, 273, 298, 323] K and RH = [20, 323]40, 60, 80, 100\\ were scanned, digitized, and fitted to low-order polynomials suitable for multilinear interpolation. For RH below 10%, a default minimum nucleation rate of 10^{-7} cm⁻³ s⁻¹ was used. The expression used for the BHNV nucleation rate, J_{BHNV} , was that presented by Vehkamäki et al. [2002], with stated range of validity T =230–305 K (and extrapolation possible down to 190K), RH from 0.01 to 100%, and $[H_2SO_4]$ from 10^4-10^{11} cm⁻³. Vehkamäki et al. [2002] stated that for $J_{\rm BHNV}$ from $10^{-6} 10^7$ cm⁻³ s⁻¹ the parameterization yields rates that are within a factor of 3 of those obtained from the full model.
- [21] The parameterization used for the ternary homogeneous nucleation rate, $J_{\rm THN}$, was that of Napari et al. [2002], who stated that over the range of temperatures 240–300 K, relative humidities 5–95%, [H₂SO₄] from 10^4-10^9 cm⁻³, [NH₃] from 0.1 to 100 ppt (parts per trillion, or pmol per mol of dry air), and $J_{\rm THN}$ from $10^{-5}-10^6$ cm⁻³ s⁻¹, the parameterized rates are within 1 order of magnitude of those given by the full theory, with closest agreement at higher values of $J_{\rm THN}$. According to Napari et al. [2002], classical nucleation theory fails near the binary H₂O-H₂SO₄ limit; therefore, in applying this parameterization here $J_{\rm BHNV}$ was used when [NH₃] < 0.1 ppt. Additionally, when [NH₃] > 100 ppt or [H₂SO₄] > 10^9 cm⁻³, the value of $J_{\rm THN}$ was set equal to that for [NH₃] = 100 ppt and [H₂SO₄] = 10^9 cm⁻³. For values of temperature or RH outside of the range of validity, $J_{\rm THN}$ was set equal to 10^{-5} cm⁻³ s⁻¹, and for $J_{\rm THN}$ > 10^6 cm⁻³ s⁻¹ it was arbitrarily set to 10^7 cm⁻³ s⁻¹.
- [22] The parameterization used for the rate of nucleation by ion-ion recombination, J_{NIIR} , was that presented by *Turco et al.* [1998]:

$$J_{\text{NIIR}} = Q_{i} f_{i} \left(\frac{[H_{2}SO_{4}]}{[H_{2}SO_{4}]_{0}} \right)^{n*},$$
 (1)

where Q_i is the local ionization (and recombination) rate, $f_i = 0.001$ is the fraction of recombination events that result in a stable nucleus at a reference vapor concentration $[H_2SO_4]_0 = 5 \times 10^6 \text{ cm}^{-3}$, and $n^* = 3$ is the threshold number of sulfuric acid molecules needed to form a stable embryo. It is further required that J_{NIIR} not exceed the ionization rate Q_i . The formulation used for Q_i is that of Yu

[2002], based on measurements of *Millikan et al.* [1944] and *Neher* [1971], which yields an increase from 2 cm $^{-3}$ s $^{-1}$ near Earth's surface to about 30 cm $^{-3}$ s $^{-1}$ at 12 km (near the height of the tropopause), with a more gradual increase with further increase in height.

[23] The nucleation rate J is an upper bound to the rate of new particle formation $J_{\rm NPF}$, as not all nuclei survive to become new particles, some being scavenged by coagulation on larger particles or on other nuclei before they grow (by condensation and by coagulation with other nuclei) to form detectable new particles, as noted in the introduction. These rates are related by

$$J_{\text{NPF}} = F_{\text{KK}}J,\tag{2}$$

where $F_{\rm KK}$ (≤ 1) is the fraction of initially formed nuclei that grow to become new particles with diameter 3 nm. An analytic formula for $F_{\rm KK}$ was presented by *Kerminen and Kulmala* [2002] and is used to calculate rates of NPF according to the nucleation parameterizations BHNJ, BHNV, THN, and NIIR. Although the derivation of their expression for $F_{\rm KK}$ is based on several assumptions, especially that self-coagulation of nuclei that are too small to be considered new particles is neglected and that the growth rate of these nuclei (here assumed to occur only through uptake of sulfuric acid, ammonia, or water) is constant, *Kerminen and Kulmala* [2002] stated that this parameterization can be successfully applied provided the nuclei number concentration remains less than 10^5-10^6 cm⁻³.

[24] An empirically derived expression for $J_{\rm NPF}$ was presented by *Eisele and McMurry* [1997] (hereinafter referred to as EM97), based on measurements of rates of NPF at Mauna Lao Observatory, Hawaii, and Idaho Hill, Colorado, reported by *Weber et al.* [1996]:

$$J_{\text{EM97}} = K[H_2 \text{SO}_4]^p. \tag{3}$$

A lower bound to the data was given by p = 1, and an upper bound by p = 2, the latter implying that the nucleation process may be collision-controlled rather than evaporation/condensation-controlled. For the present calculations p = 2 and $K = 3.5 \times 10^{-15}$ cm³ s⁻¹, the mean value of those corresponding to the fits shown in EM97 for p = 2. As the accuracy of such an empirical expression can be expected to vary considerably, both spatially and temporally, a single set of values (p, K) would not be expected to be appropriate for a region as large as the continental United States; hence the rates obtained from this expression must be treated as representative of potential rates.

- [25] The dependence of $J_{\rm NPF}$ on [H₂SO₄] is shown in Figure 1 for the five NPF parameterizations BHNJ, BHNV, THN, NIIR (each with the factor $F_{\rm KK}$), and EM97, at 298 K, 50% RH, 10 and 100 ppt of [NH₃], and with no background aerosol and with substantial background aerosol (a lognormal size distribution with $N_{\rm tot}=500~{\rm cm}^{-3}$, geometric standard deviation 1.5, and geometric mean diameter 150 nm). For each NPF parameterization (except for EM97) a minimum background value of $J_{\rm NPF}$ is arbitrarily taken as $1\times10^{-7}~{\rm cm}^{-3}~{\rm s}^{-1}$, and a maximum value is arbitrarily taken as $1\times10^{7}~{\rm cm}^{-3}~{\rm s}^{-1}$.
- [26] The value of $[H_2SO_4]$ that results in a given J_{NPF} depends strongly on the NPF formulation (and on the amount of background aerosol; because H_2SO_4 is scavenged by

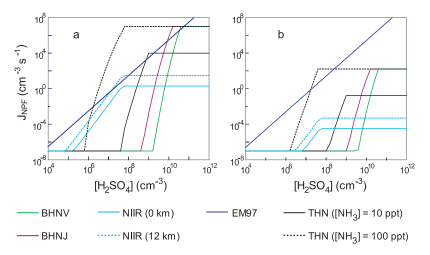


Figure 1. New particle formation rate $J_{\rm NPF}$ versus sulfuric acid concentration [H₂SO₄] for the several new particle formation parameterizations considered here at 25°C and relative humidity 50%: (a) without background aerosols and (b) with high concentration of background aerosols. A lognormal with $N_{\rm tot} = 500~{\rm cm}^{-3}$, $\sigma_{\rm g} = 1.5$, and $D_{\rm g} = 150~{\rm nm}$ is used, corresponding to a rate constant for condensational sink for sulfuric acid of $k_{\rm cs} = 2 \times 10^{-2}~{\rm s}^{-1}$. The effect of background aerosols on $J_{\rm NPF}$ was not considered for $J_{\rm NPF}$ of *Eisele and McMurry* [1997] (EM97). BHNV, binary homogeneous nucleation parameterization of *Vehkamäki et al.* [2002]; BHNJ, that of *Jaecker-Voirol and Mirabel* [1989]; NIIR, nucleation by ion-ion recombination; THN, ternary homogeneous nucleation.

existing aerosol, NPF is suppressed at high aerosol concentration). Typically this value is the least for EM97 and THN (for $[NH_3] = 100$ ppt), slightly greater for NIIR, several orders of magnitude greater for THN (for $[NH_3] = 10$ ppt), and greater still for BHNJ and BHNV. As the dependence of J_{NPF} on $[H_2SO_4]$ is much stronger for THN (for $[NH_3] = 100$ ppt) than for EM97, when sufficient ammonia is present THN is the dominant nucleation mechanism at values of $[H_2SO_4]$ sufficiently great that appreciable NPF occurs.

2.2. Treatment of Sulfuric Acid

[27] Sulfuric acid is formed by reaction of OH with SO₂, which, in the presence of water vapor and oxygen, occurs by the overall reaction

(R1)
$$SO_2 + OH \rightarrow ... \rightarrow H_2SO_4$$

with rate $k_1[OH][SO_2]$, where k_1 is a temperature-dependent constant. Sulfuric acid vapor is removed by NPF and by condensation on existing aerosol particles, the latter process occurring at a rate $(-d[H_2SO_4]/dt)_{cond} = k_{CS}[H_2SO_4]$ with effective first-order rate constant k_{CS} , which depends primarily on the concentrations of surface area in the Aitken and accumulation modes.

[28] In the conventional operator-splitting approach as employed in CMAQ, the sulfuric acid concentration calculated in the chemistry module is introduced into the aerosol nucleation and dynamics modules, where it is partitioned between nucleation and condensation. The approach currently employed in CMAQ assumes that [H₂SO₄] is in steady state with condensation being the only sink:

$$\frac{d[H_2SO_4]}{dt} = 0 = k_1[OH][SO_2] - k_{CS}[H_2SO_4], \tag{4}$$

where the first term on the right side represents the average production rate of sulfuric acid molecules over the time interval. When this approach was employed with NPF from both THN and NIIR, it was found that $J_{\rm NPF}$ could be erroneously high and that the amount of sulfuric acid consumed by NPF could become comparable to or exceed the rate of condensation on existing particles, or could even exceed the amount formed. An attempt was made to circumvent this problem by invoking a mass cap: a limit on the amount of $\rm H_2SO_4$ that can go into new particle formation so that it does not exceed the amount formed.

[29] The probability distribution function of $J_{\rm NPF}$ according to this approach, $dF/d\log J_{\rm NPF}$, the fraction of occurrences over the entire modeling domain and period that are within a given logarithmic range of $J_{\rm NPF}$, is shown in Figure 2a both without and with imposition of the mass cap (the total possible number of occurrences is equal to the number of grid cells in the domain, $\sim 1.2 \times 10^5$, times the number of time intervals in the model run, $\sim 3 \times 10^3$, i.e., $\sim 4 \times 10^8$). The imposition of the mass cap greatly reduces the number of instances of high values of $J_{\rm NPF}$ (Figure 2a).

[30] To account for the sink of sulfuric acid by NPF, a second approach was employed in which, at steady state, the rate of production of sulfuric acid is set equal to its rate of loss by condensation and new particle formation:

$$\frac{d[H_2SO_4]}{dt} = 0 = k_1[OH][SO_2] - k_{CS}[H_2SO_4] - nJ_{NPF}([H_2SO_4]),$$
(5)

where the dependence of $J_{\rm NPF}$ on the sulfuric acid concentration is explicitly shown and n, the number of sulfuric acid molecules in a new particle, is taken as 128 under the assumption that each new particle of diameter 3 nm consists mainly of sulfuric acid. This expression yields

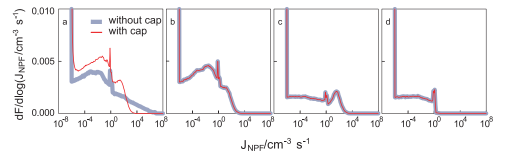


Figure 2. Probability distribution function of $J_{\rm NPF}$, $dF/d\log J_{\rm NPF}$, the fraction of occurrences over the entire modeling domain and period that are within a given logarithmic range of $J_{\rm NPF}$, as a function of new particle formation rate $J_{\rm NPF}$: (a) calculated with THN+NIIR using the standard Community Multiscale Air Quality (CMAQ) operator-splitting algorithm, (b) calculated with THN+NIIR using the steady state $H_2{\rm SO}_4$ concentration obtained for new particle formation and condensation occurring simultaneously, (c) calculated for THN+NIIR without the steady state assumption for $H_2{\rm SO}_4$, and (d) calculated for NIIR acting alone without the steady state assumption for $H_2{\rm SO}_4$. Red denotes calculations with "mass cap" which limits $J_{\rm NPF}$ to the amount of $H_2{\rm SO}_4$ formed in the time interval; gray denotes calculations without the mass cap.

a parametric expression for steady state sulfuric acid concentration:

$$[H_2SO_4] = \frac{k_1[OH][SO_2] - nJ_{NPF}([H_2SO_4])}{k_{CS}},$$
 (6)

which can be solved iteratively for a given formulation of $J_{\rm NPF}$. Loss of $H_2{\rm SO}_4$ was taken into account by initially calculating the rates of consumption of $H_2{\rm SO}_4$ by NPF and condensation independently, as if all of the available $H_2{\rm SO}_4$ could be consumed by either process alone, and then, if the total $H_2{\rm SO}_4$ consumed by the two processes exceeded the total amount available, $[H_2{\rm SO}_4]$ was decreased, resulting in a decrease in both NPF and condensation rates, until the total available $H_2{\rm SO}_4$ was consumed. The problem of overconsumption of $[H_2{\rm SO}_4]$ was resolved by this approach, as shown by the lack of noticeable change in the probability distribution due to imposition of the mass cap in Figure 2b.

[31] In some circumstances, however, the steady state assumption is not valid, as when the time constant for approaching steady state is comparable to or greater than the time interval used in the model. Under these circumstances, a third approach was employed in which $[H_2SO_4]$ was evaluated as the average of $[H_2SO_4]$ at the start and the end of a time interval, calculated from

$$\frac{d[H_2SO_4]}{dt} = k_1[H][SO_2] - k_{CS}[H_2SO_4] - nJ_{NPF}([H_2SO_4])$$
 (7)

rather than by using the steady state assumption. When this third approach was applied the probability distribution (Figure 2c) exhibited further substantial suppression of NPF relative to that calculated under the steady state assumption (Figure 2b). This approach to taking into account the time dependence seemed to yield reasonable results and to work over the entire range of $J_{\rm NPF}$ considered, and thus was selected for the present study. Additionally, as $J_{\rm NPF}$ is held constant over each 15 min time interval, the possibility of a more rapid pulse nucleation event is precluded.

[32] In summary, when steady state conditions, in which the rate of production of [H₂SO₄] is nearly balanced by its

loss to condensation and to new particle formation, are rapidly (i.e., within a time step) approached, equation (6) is used; otherwise equation (7) is solved to find the sulfuric acid concentration. In neither situation is a mass cap employed. It should be noted that NPF is a rare phenomenon, with considerably less than 1% of occurrences exhibiting values of $J_{\rm NPF}$ greater than 1 \times 10⁻⁶ cm⁻³ s⁻¹, and even fewer greater than 1 cm⁻³ s⁻¹ (Figure 2c), a value typically taken as that which would produce an appreciable number of new particles. One additional feature visible in each approach is a peak in $J_{\rm NPF}$ near 1 cm⁻³ s⁻¹; this results from NIIR, as is evidenced by the occurrence of this peak when the third approach is applied to nucleation from NIIR alone (Figure 2d).

2.3. Evolution of Particle Size

[33] Evolution of particle size in each mode is represented in this model, as in CMAQ, by the rate of change with time of three moments of the size distribution of number concentration with respect to the dry particle diameter: the zeroth moment (which is the same as the number concentration N), the second moment M_2 , (directly proportional to the surface area concentration A), and the third moment M_3 (directly proportional to the volume concentration V). Each moment in each mode is transported, preserving the lognormal distribution, as in the work of Binkowski [1999]. As the size distribution of number concentration in each mode is assumed to be a lognormal, the geometric mean dry diameter D_g and the geometric standard deviation σ_g of each mode are related to these moments by

$$D_g = \frac{M_2^{3/2}}{N^{5/6} M_3^{2/3}},\tag{8}$$

and

$$\ln \sigma_g = \left(\frac{\ln N + 2\ln M_3 - 3\ln M_2}{3}\right)^{1/2} = \left[\ln\left(\frac{N^{1/3}M_3^{2/3}}{M_2}\right)\right]^{1/2},$$
(9)

respectively. Major processes affecting the size distribution affect the moments, and hence $D_{\rm g}$ and $\sigma_{\rm g}$, in different ways. Coagulation of particles within the same mode reduces Nand M_2 but leaves M_3 unchanged; because in this process smaller particles attach to larger ones within the same mode, it results in an increase in $D_{\rm g}$ and a decrease in $\sigma_{\rm g}$. Condensation of gases (other than water vapor) increases M_2 and M_3 but leaves N unchanged, and because smaller particles experience greater relative growth than larger ones, this process also typically results in an increase in $D_{\rm g}$ and a decrease in σ_g . NPF increases all three moments of the Aitken mode, but especially N, and as the particles that are produced are very small ones, this process results in a decrease in $D_{\rm g}$ and an increase in $\sigma_{\rm g}$ in that mode. Condensation and evaporation of water vapor are taken into account as described above, and particle diameters used in calculations are "wet" diameters, meaning that they are the physical sizes of particles in equilibrium at the ambient relative humidity; however, the diameters and moments are reported as "dry," meaning that water mass is not included.

[34] Intermodal transfer (IMT) is an artificial process that prevents the Aitken mode from merging into the accumulation mode by transferring some fraction of each of the three moments of the Aitken mode to the accumulation mode at the end of each time interval when certain requirements are fulfilled, namely that both N and the rate of condensational growth of M_3 in the Aitken mode are greater than those in the accumulation mode [Binkowski and Roselle, 2003]. This transfer results in a decrease in all three moments in the Aitken mode and an increase of the three moments in the accumulation mode. Because the particles that are transferred are the largest particles in the Aitken mode, and among the smallest in the accumulation mode, IMT results in decreases in D_{g} of both the Aitken mode and the accumulation mode. The fractions of the moments in the Aitken mode resulting from particles with diameters greater than that for which the corresponding concentrations of both modes are equal are transferred to the accumulation mode with the added imposition that not more than one half of the third moment of the Aitken mode can be transferred at any one time step. As the size distributions of the concentrations in each mode are assumed to be lognormals, the diameters at intersection are readily calculated by solving quadratic equations.

2.4. Input Data to the Regional Model Calculations

[35] Meteorological data used for the chemical transport model calculations (T, RH, wind speed and direction, cloud and rain properties, and turbulent fluxes) were generated from the Mesoscale Model version 5 (MM5) [Dudhia, 1993] with the four-dimensional data assimilation option (which forces the output to conform to observations) on an hourly basis using NCEP (National Centers for Environmental Prediction) reanalysis for the continental United States with horizontal resolution of 60 km for the continental-scale domain and 12 km for the northeastern United States. These data were then input into the Meteorology-Chemistry Interface Processor (MCIP) [Byun et al., 1999], which links MM5 to CMAQ in terms of data format, units, and reconstruction on a different grid structure. MCIP also enforces consistency among the meteorological variables such as height of the planetary boundary layer and cloud parameters.

- [36] Anthropogenic emissions (gas and particulate) were based on the Environmental Protection Agency (EPA) 1999 National Emission Inventory (NEI99) updated for a typical summer day of 2004 using projected and known changes for the previous 5 years according to McKeen et al. [2005]. In particular, Canadian area sources are included south of 52°N but Canadian point sources are not included in this inventory. This revised inventory was previously used for evaluation of air quality forecast models in the ICARTT field study [Fehsenfeld et al., 2006]. The primary emitted species are NO, NO₂, VOC, CO, SO₂, NH₃, PM2.5, and PM10, along with 41 speciated VOC compounds and 5 PM2.5 aerosol species (sulfate, nitrate, organic carbon, elemental carbon, and unspecified PM2.5). Dimethylsulfide (DMS), a marine indirect source of SO₂ through its oxidation, was not treated in this study, thus precluding any potential contribution to [H₂SO₄] (through oxidation of SO₂) and hence nucleation from this source. The inventory was transformed by the Sparse Matrix Operator Kernel Emissions (SMOKE) modeling system (available at http:// www.baronams.com/products/smoke/) to hourly gridded emission rates at a resolution of 4 km, which were remapped to the grid sizes employed in the two model domains (60 km and 12 km). Area sources were assigned to the lowest vertical level and point sources to specific vertical layers depending on quantities such as stack height, plume rise, and vertical dispersion. Emissions that are introduced in a single grid cell in the model are treated as uniformly distributed over the cell.
- [37] The emission inventory for primary particles contains no information on their sizes. In the current study the size distributions of emitted mass are treated as lognormals with dry geometric mean diameters of 30 nm, 300 nm, and 6 μ m and geometric standard deviations of 1.7, 2.0, and 2.2 for the Aitken, accumulation, and coarse modes, respectively. The corresponding size distributions of aerosol number emissions in these three modes are therefore also lognormals with dry geometric mean diameters near 13, 71, and 930 nm, respectively, and the same geometric standard deviations as for the mass emissions. The consequences of these assumptions, especially for geometric mean diameters, are examined below. The partitioning of each species between modes follows the CMAQ approach of putting 99.9% of total emitted mass into the accumulation mode and 0.1% into the Aitken mode, both in the dry state (this is equivalent to assuming that nearly twice as many particles are emitted into the accumulation mode as into the Aitken mode); the consequences of this choice are also discussed below. The coarse mode emissions are based on the difference of PM10 and PM2.5. No explicit emission of dust or sea salt particles, which would be mainly in the coarse mode, is treated in the present model; thus the condensation sink in the model might be less than what it would be in situations and locations for which these particles would otherwise contribute.
- [38] Rates of biogenic emissions of gases were obtained from the Biogenic Emissions Inventory System (BEIS) version 3.11, developed by the U.S. Environmental Protection Agency (described at http://www.epa.gov/asmdnerl/biogen.html). Emissions of isoprene, monoterpenes, other organic VOC, and NO were partitioned into the appropriate SAPRC-99 species according to meteorological conditions.

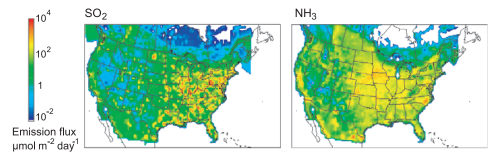


Figure 3. Spatial distribution of primary emission fluxes for new particle formation (NPF) precursors, SO₂ and NH₃, for 1 July 2004.

Emissions from biomass burning were not included in source inventories.

[39] The spatial distributions of emission fluxes of sulfur dioxide and ammonia, which are precursors for NPF, are shown in Figure 3 for 1 July 2004. Emission fluxes of sulfur dioxide were high over the eastern United States, mainly because of the density of power plants there. In contrast, emission fluxes of ammonia were high over the midcontinental United States, where agriculture activities and livestock husbandry are centered.

3. Experiments

[40] The sensitivities of aerosol number concentration to NPF and primary emission were examined using nine model variants as listed in Table 1. Sensitivity to NPF formulation was examined by comparing the base case (denoted BASE), for which the sole source of particles was primary emission (i.e., no NPF), to two BHN formulations (BHNJ and BHNV), one THN formulation, and NIIR (for all of which the conversion factor F_{KK} was employed to yield J_{NPF}), and one empirically derived parameterization for $J_{\rm NPF}$ (EM97), each of which was examined in a separate model run. Sensitivity to the number (size) of emitted particles was examined by comparing results for the NPF mechanism being the sum of THN and NIIR with the emissions as described above (D_0 case) to those for which the dry diameters of the emitted particles were decreased and increased by a factor of two (D/2 and $D\times 2$ cases, respectively) while holding the mass emissions constant. In each case the model was run for 2 months, the first with no nucleation and the second with nucleation, before the time period of interest. The model was initialized using a typical profile of chemical species provided by CMAQ (thus no

Table 1. Model Variants and Their Descriptions

Model Variant	Nucleation Formulation	$\frac{D_{\rm g,emission}}{D_{\rm g,emission,BASE}}$	
BASE	none	1	
BHNV	BHN [Vehkamäki et al., 2002]	1	
BHNJ	BHN [Jaecker-Voirol and Mirabel, 1989]	1	
NIIR	NIIR [Turco et al., 1998]	1	
EM97	empirical NPF parameterization	1	
	[Eisele and McMurry, 1997]		
THN	THN [Napari et al., 2002]	1	
D_0	THN + NIIR	1	
D/2	THN + NIIR	0.5	
$D\times 2$	THN + NIIR	2.	

intercontinental transport of pollutants nor any downward transport of ozone from the stratosphere was included).

3.1. Sensitivity to NPF Formulation

[41] The dependence of aerosol number concentration and properties on NPF formulation was investigated by analyzing in detail model results at four sites, chosen to represent typical values for different parameters that might affect NPF. These sites are shown in Figure 4 along with the simulated wind field, averaged over July 2004 at an elevation corresponding to 85% of the surface pressure (near 1.5 km). Site A, near the northwestern corner of Iowa, is a rural location surrounded by intensely fertilized agriculture and is characterized by high [NH₃]. Site B, near New Haven, Connecticut, is an urban site that is strongly impacted by upwind urban emissions. Site C, near Acadia National Park, Maine, is a location that is frequently subject to transport of material from urban locations, although further removed than Site B. Site D, 350 km southeast of Cape Cod, Massachusetts, is a marine location well removed from local continental sources. Time series of $J_{\rm NPF}$ and controlling variables (T, RH, [SO₂], [NH₃], and [H₂SO₄]), number concentrations in the Aitken (N_{ATK}) and accumulation (N_{ACC}) modes, and other related quanti-

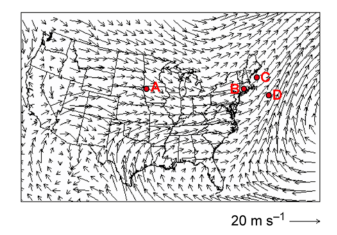


Figure 4. Model domain with simulated wind field, averaged over July 2004, at an elevation corresponding to 85% of the surface pressure. Red dots denote the sensitivity analysis sites: A, northwestern Iowa (96.2°W, 43.5°N); B, near New Haven, Connecticut (72.9°W, 41.1°N); C, near Acadia National Park, Maine (68.1°W, 43.7°N); D, 350 km southeast of Cape Cod, Massachusetts (66.5°W, 39.9°N).

ties for each NPF formulation at the lowest model level (0–20 m above the ground) at each of the four locations are shown in Figures 5a–5d. As noted above, values of $D_{\rm g,ATK}$, $D_{\rm g,ACC}$, and the volume concentration in the Aitken mode $V_{\rm ATK}$ are reported for the dry particles (i.e., without taking into account the associated water), although in the calculation of processes such as dry deposition and coagulation the actual (i.e., wet) sizes, which included the associated water, were used.

[42] The behavior of factors affecting NPF differed appreciably between the sites. Temperatures at Sites A and B exhibited considerable variability and had pronounced diurnal cycles, as opposed to those at Sites C (which were consistently lower than the others) and D. Relative humidities at Sites A and B likewise exhibited considerable variability and had pronounced diurnal cycles, whereas those at Sites C and D were consistently higher and much more uniform, reflecting the coastal or marine locations of these sites. The concentrations of SO₂ at Site B were considerably higher than those at the other sites (note the different axis scale for [SO₂] for that site), whereas those at Sites C and D were often negligible, with only occasional instances of nonnegligible values. The concentrations of NH₃ at Site A were considerably higher than those at the other sites (note also the different axis scales for [NH₃] for each site), which were frequently negligible. Thus it was expected that NPF at these sites by the several formulations examined here would exhibit substantial differences, both in frequency and amount, allowing differences in controlling processes to be identified and their sensitivities examined.

[43] Several features were common among all sites. The concentration of sulfuric acid exhibited a strong diurnal profile, generally peaking around local noon, reflecting the photochemical production of OH and the short lifetime of this species. As NPF requires the presence of H₂SO₄, both $F_{\rm KK}$ and $J_{\rm NPF}$ for each of the formulations (when NPF occurred) also exhibited a diurnal profile, but with the magnitude of J_{NPF} , or whether NPF occurred at all, depending on location and, at a given location, on the NPF formulation. For instance, because THN is governed by the availability of NH₃, NPF by THN occurred mainly in regions with appreciable [NH₃]. As NPF results in new particles in the Aitken mode, NATK likewise exhibited a strong diurnal pattern when NPF occurred, but with a somewhat asymmetric or saw-tooth profile reflecting the abrupt onset of NPF followed by a slower decay of N_{ATK} due to coagulation and/or intermodal transfer. Self-coagulation of Aitken mode particles also exhibited a diurnal

profile, but with the maxima occurring slightly after those for $J_{\rm NPF}$.

[44] The number concentration in the Aitken mode was generally considerably greater than that in the accumulation mode, whereas the concentrations of surface area and volume (and thus mass) in the accumulation mode were considerably greater than those in the Aitken mode. $V_{\rm ATK}$ exhibited much less variation among NPF formulations than did $N_{\rm ATK}$, as new particles, because of their size, contribute little to volume concentrations in that mode. $N_{\rm ACC}$ was nearly independent of NPF formulation in most instances, implying that its major source was emissions. The main removal process of $N_{\rm ACC}$ was wet deposition, as shown by sharp decreases in $N_{\rm ACC}$ coincident with precipitation events.

[45] For each NPF formulation $D_{g,ATK}$ typically exhibited a diurnal variation opposite of that of J_{NPF} (Figure 5), decreasing as new 3-nm particles were formed during the daytime and increasing because of coagulation at night. When appreciable NPF occurred, $D_{g,ATK}$ at the lowest model level was the smallest (a few nanometers) at local noon and the largest (several tens of nanometers) at night. The lowest daytime $D_{g,ATK}$ generally occurred for THN but the greatest occurred, unexpectedly, for BHNJ. This was attributed to NPF occurring at higher altitudes (compare Figure 6), with new particles being rapidly transported to the surface where they underwent growth by condensation and coagulation. This inference is supported by the finding that large values of $D_{\rm g,ATK}$ were accompanied by high self-coagulation rates and high sulfate concentrations in the Aitken mode, for example, on 3 July at Site A (Figure 5a). In contrast to $D_{g,ATK}$, there was very little temporal variation of $D_{g,ACC}$, this quantity being near 70 nm (corresponding to the geometric mean dry diameter of the number size distribution of emissions, as noted above) at each site, independent of NPF formulation, except for abrupt decreases coincident with precipitation events (shown by occurrences of high wet deposition), in which larger particles are preferentially activated to form cloud drops and subsequently removed. This finding also suggests that the dominant source of accumulation mode particles was primary emissions.

[46] When IMT occurred, it exhibited diurnal variation similar to that of $J_{\rm NPF}$. However, the occurrence of IMT at the several sites was quite intermittent, being associated with NPF (no IMT occurred for the BASE case) but not occurring in all instances of NPF. The dependence of IMT on $J_{\rm NPF}$ and $[{\rm H}_2{\rm SO}_4]$ is examined in Figure 7, which shows IMT fraction, the fraction of time intervals for which IMT

Figure 5a. Time series of T, RH, [SO₂], [NH₃], [H₂SO₄], $J_{\rm NPF}$, $F_{\rm KK}$, $N_{\rm ATK}$, $V_{\rm ATK}$, $N_{\rm ACC}$, $D_{\rm g,ATK}$, $D_{\rm g,ACC}$, rate of intermodal transfer (IMT) of aerosol number, intramodal coagulation rates for the Aitken and the accumulation modes, dryair molar mixing ratios of sulfate and ammonium in the Aitken and accumulation modes, mass concentration of secondary organic aerosol (SOA) in the Aitken mode, wet deposition flux of accumulation mode particle number, and dry deposition flux of Aitken mode particle number at the lowest level (0–20 m above the ground) for different model variants at Site A during July 2004. Tick marks denote midnight UTC; dashed vertical blue lines denote local noon. THN (black line) is plotted last and may obscure traces of other model variants. The quantities $D_{\rm g,ATK}$, $D_{\rm g,ACC}$, and $V_{\rm ATK}$ are reported for dry particles, although for use in evaluation of coagulation rates, etc., they were calculated for particles at their ambient (i.e., wet) sizes. Abundances of [SO₂], [NH₃], [SO₄²⁻], and [NH₄⁺] are presented as molar mixing ratios in dry air, nmol mol⁻¹ (ppb, parts per billion). At 298 K and 1 atm, a molar mixing ratio of 1 ppb is equivalent to a mass concentration of 3.9 μ g m⁻³ for ammonium.

occurred anywhere in the columns over the four sites (i.e., including all vertical layers) for the entire month of July. For low values of $J_{\rm NPF}$ (i.e., <10 cm⁻³ s⁻¹), IMT fraction is quite low (<15%) and exhibits a weak dependence on $J_{\rm NPF}$, but this fraction increases strongly with increasing $J_{\rm NPF}$ for larger values, approaching 100% for $J_{\rm NPF}=10^3$ cm⁻³ s⁻¹ (Figure 7a). In contrast, IMT fraction is nearly independent

of [H₂SO₄] above a minimum value for J_{NPF} in the range $10-100 \text{ cm}^{-3} \text{ s}^{-1}$ (Figure 7b).

[47] Aitken and accumulation mode aerosol particles were typically composed mainly of sulfate and ammonium, the acidity of particles being determined by the availability of ammonia, which differed among the sites. In some instances there was an appreciable contribution from sec-

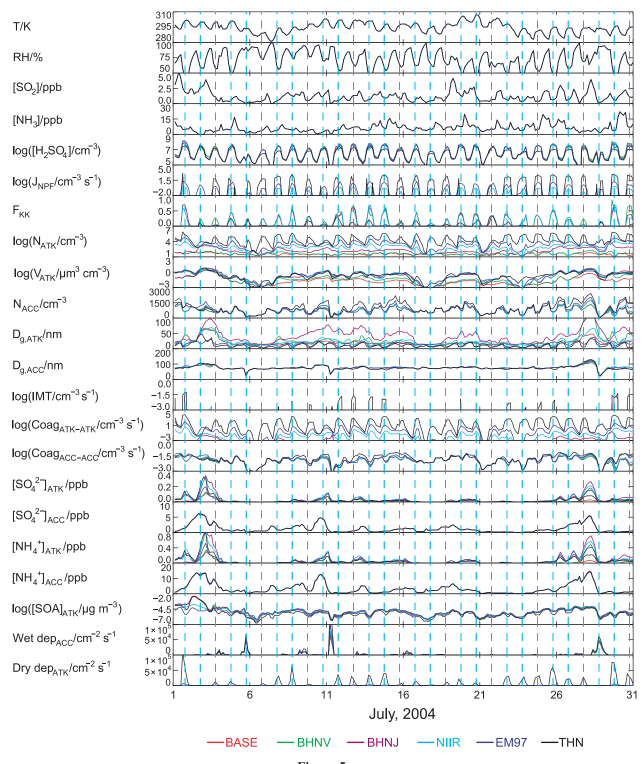


Figure 5a

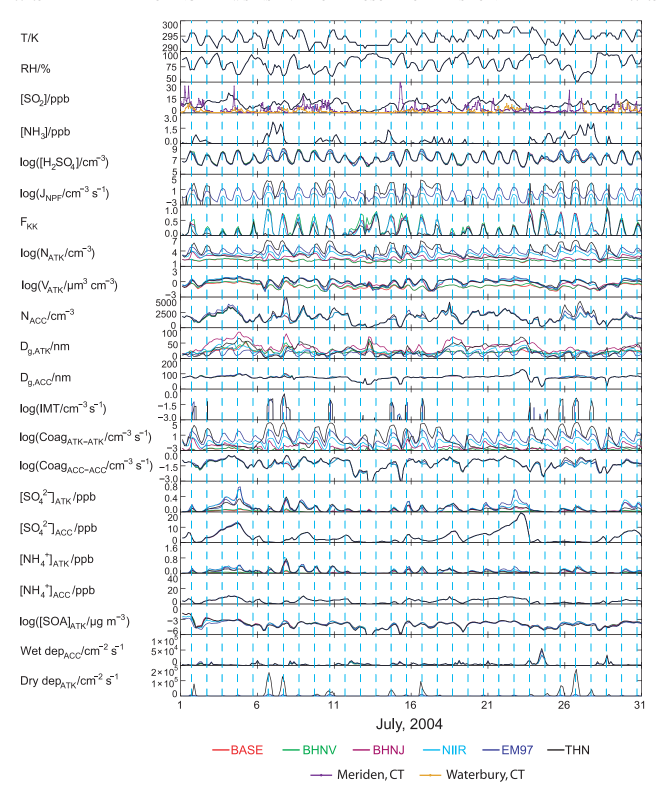


Figure 5b. Same as Figure 5a except for Site B, with SO_2 mixing ratio measurements from Environmental Protection Agency monitoring stations at Meriden and Waterbury, Connecticut. Note the different scale for some quantities such as $[SO_2]$.

ondary organic aerosol, SOA, although this typically comprised at most a few percent of total particle mass in the Aitken and accumulation modes. SOA is reported as often constituting a substantial fraction of the mass of fine

particles in the atmosphere [Saxena and Hildemann, 1996; Zhang et al., 2007], which would imply that the contribution of SOA to particle growth may be underestimated in the

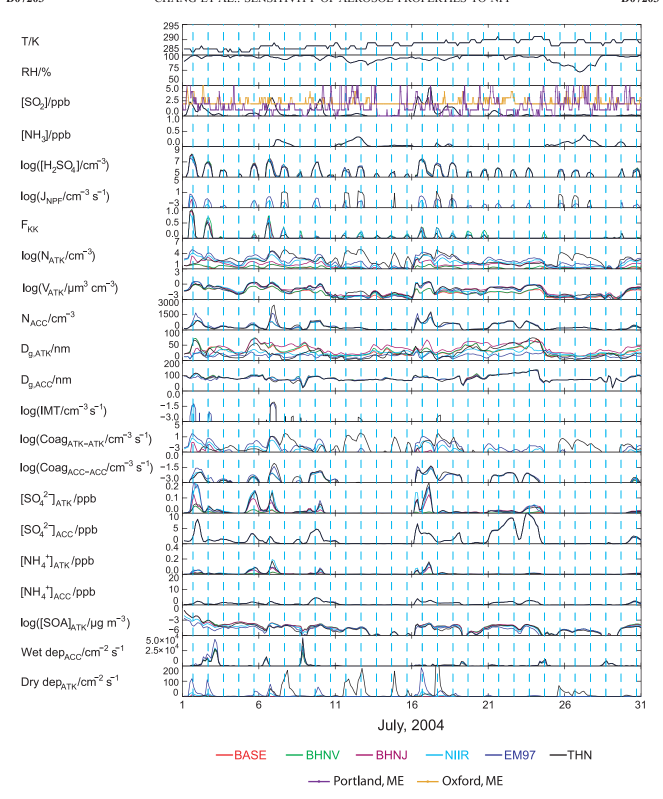


Figure 5c. Same as Figure 5a except for Site C, with SO₂ mixing ratio measurements from Environmental Protection Agency monitoring stations at Oxford and Portland, Maine.

present model. However, as discussed below, this does not seem to always be the situation.

[48] As expected, there were considerable differences in NPF between the sites. At Site A, a rural location, $[H_2SO_4]$ frequently exceeded 10^7 cm⁻³ near local noon and $[NH_3]$

was typically quite high. Under these conditions there were large differences in the frequency and rate of NPF between the different formulations, with THN exhibiting the highest rates of NPF, followed by EM97 and NIIR. The fraction of time intervals during which NPF occurred in the lowest

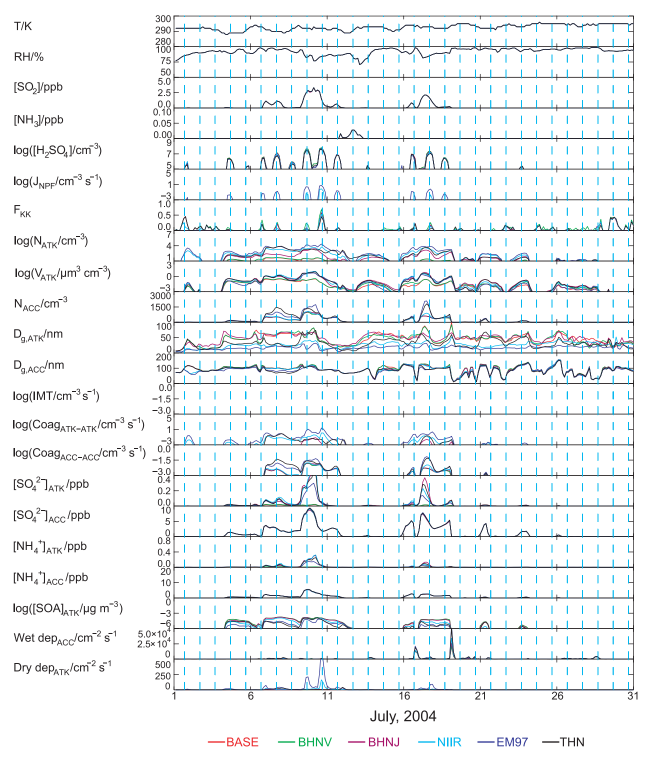


Figure 5d. Same as Figure 5a except for Site D.

model level at rates greater than $1 \text{ cm}^{-3} \text{ s}^{-1}$ was greatest for THN with 37% (out of 248 time intervals), followed by 29% for EM97 and only 1.6% for NIIR. NPF did not occur at all for BHNJ or BHNV, as the modeled $[\text{H}_2\text{SO}_4]$ was insufficient for NPF at the high temperatures at this location. THN frequently yielded rates of NPF that exceeded those typical of urban areas, $\sim 100 \text{ cm}^{-3} \text{ s}^{-1}$ [Kulmala et al., 2004], and because the modeled [NH₃] exceeded the parameterization limit given by Napari et al. [2002] of

100 ppt (above which the modeled $J_{\rm NPF}$ was held equal to the maximum given by the parameterization), the actual values of J and $J_{\rm NPF}$ may have even exceeded the values shown. These considerations suggest that $J_{\rm NPF}$ according to THN exhibits high uncertainty in regions of high [NH₃]. Although BHNJ resulted in no NPF at the lowest model level, it yielded greater $N_{\rm ATK}$ than the BASE case did, suggesting rapid transport of new particles from above. The fraction of time intervals for which IMT occurred at the

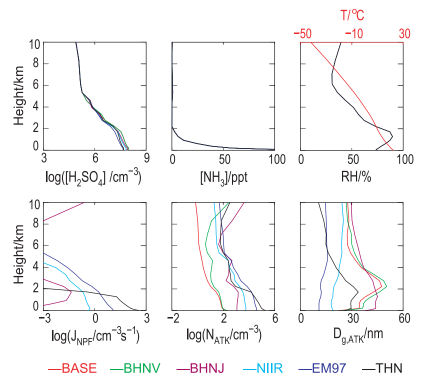


Figure 6. Vertical profiles of $[H_2SO_4]$, $[NH_3]$, T and RH, J_{NPF} , N_{ATK} , and $D_{g,ATK}$ at local noon for different model variants, averaged over July 2004 at Site B. Abundance of $[NH_3]$ is presented as molar mixing ratio in dry air, pmol mol⁻¹ (ppt, parts per trillion).

lowest level at Site A was 18% for THN but only 2.4% for EM97. NPF without IMT can occur because of the requirement for IMT that the condensational growth rate of aerosol mass in the Aitken mode must be greater than that in accumulation mode.

[49] The concentrations of (dry) particle volume in the Aitken mode $V_{\rm ATK}$ at this site for THN, EM97, NIIR, and BHNJ were roughly the same, and they were typically greater than those in BASE and BHNV by an order of magnitude because of newly formed particles, which were too small to enhance $V_{\rm ATK}$ at their initial size, but which increased $V_{\rm ATK}$ by their further growth by condensation of substances (other than water) with rates constrained by available condensing vapor.

[50] Although the time dependences of the various quantities of interest at Site B (Figure 5b), semi-urban Connecticut, were similar to those at Site A, there were noticeable differences, especially in $J_{\rm NPF}$, $N_{\rm ACC}$, the occurrence of IMT, and $D_{\rm g,ATK}$. [H₂SO₄] was frequently greater than at Site A, and [NH₃] was typically considerably lower because of the molar mixing ratio of sulfate commonly exceeding half that of ammonium, resulting in unneutralized acid, for example, 3–5 July. Formulations exhibiting the greatest $J_{\rm NPF}$ and $N_{\rm ATK}$ were either THN, for which NPF was intermittent, or EM97. As for Site A, BHN was not capable of NPF because the modeled [H₂SO₄] (\leq 10⁸ cm⁻³) was insufficient for NPF at this warmer location (T > 290 K). The fraction of time intervals for which IMT occurred at the lowest model level for EM97 increased from 2.4% at Site A

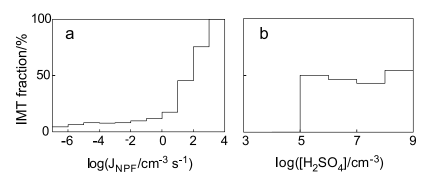


Figure 7. IMT fraction, the fraction of time intervals for which IMT occurred anywhere in the columns over the four sites shown in Figure 4 during July 2004, in percent, as a function of (a) $J_{\rm NPF}$ and (b) $[{\rm H_2SO_4}]$ for $J_{\rm NPF}$ in the range $10-100~{\rm cm}^{-3}~{\rm s}^{-1}$.

to 17% at Site B because the greater $[H_2SO_4]$ resulted in greater $J_{\rm NPF}$ and consequently more frequent IMT according to this formulation. $D_{\rm g,ATK}$ was greater for THN than for EM97, and even the greatest among all NPF formulations considered in some instances, because the intermittent nature of NPF by this formulation allowed more time for new particles to grow by self-coagulation and condensation.

[51] At Site C, the remote downwind site, [SO₂] and [NH₃] were generally considerably lower (Figure 5c) than at Sites A and B. Some instances of high values occurred, but these can be attributed to transport from different locations. Insight into the origins of SO₂ and NH₃ can be found in maps of emissions of these substances (Figure 3); strong sources of SO₂ lie south of this site, whereas strong sources of NH₃ lie both north and south of this site. During transit NH₃ would be consumed to neutralize H₂SO₄, indicating that instances of simultaneous high [H2SO4] and [NH3] would be rare at this site. In addition F_{KK} was typically low because [H₂SO₄] was quite low despite low background aerosol concentration. Under these conditions NPF events were weaker and less frequent than at Site A or B, and, when they occurred, were mainly by THN in situations of northwesterly winds. On the other hand, the number of occurrences of IMT was not less than that at Site A or B, although the magnitude of such events generally was too small ($<10^{-3}$ cm⁻³ s⁻¹) to influence $N_{\rm ACC}$ appreciably. As the occurrence of IMT does not depend strongly on $J_{\rm NPF}$ when J_{NPF} is less than 10 cm⁻³ s⁻¹ (Figure 7a), most instances of IMT at this site could not be explained by local NPF and must therefore have involved Aitken particles brought to the site by either vertical or horizontal transport.

[52] At Site D, the remote marine site, [SO₂] and [NH₃] were generally also considerably lower (Figure 5d) than at Sites A and B. [H₂SO₄] exceeded 10⁶ cm⁻³ only when the site was under the influence of transport from the adjacent continent. For example, on 10 July Site D was located under the east flank of a low-pressure system which transported SO₂ leaving the continent at Delaware along 38°N under a strong southwesterly wind. The production of H₂SO₄ from local oxidation of SO₂ advected into this region resulted in a few significant NPF events, and only by EM97. NPF by THN did not occur because ammonia had been totally consumed to neutralize acidic species before leaving the continent, and NPF by BHNV or BHNJ did not occur in the lowest model level because of the high temperature there ($T \sim 290$ K). However, events of NPF by EM97 were not accompanied by increases in concentrations of surface area or volume in the Aitken mode because of insufficient condensable vapor; consequently EM97 had the lowest $D_{g,ATK}$ among all model variants. $D_{g,ATK}$ was typically higher for the BASE case, as no IMT occurred and particles continued to grow without being removed from the Aitken mode. Omission of sea salt and DMS emissions in the model would be expected to impact this site more than any of the others. The main consequence of sea salt would have been a greater condensational sink; however, as concentrations of condensable vapor at this site were often low (Figure 5d), the consequences of not including sea salt were probably minor. DMS through its oxidation would result in SO₂, and through its oxidation H₂SO₄, suggesting that concentrations of these substances may have been underestimated, but the air mass at this location was certainly strongly continentally influenced (Figure 3) with relatively little time spent over the water for oxidation to occur, so the consequences of not including DMS are probably minor as well.

[53] An attempt was made to evaluate the representativeness of mixing ratios of SO₂ used in calculation of the H₂SO₄ production rate by comparison with measurements made at nearby U. S. Environmental Protection Agency air quality monitoring stations for the several locations examined in Figure 5. Such evaluations are particularly difficult for a primary emittant such as SO₂, which derives mainly from point sources and which consequently exhibits considerable short-term and subgrid variability relative to calculations obtained with a continental-scale model such as employed here. A further limitation on such comparisons is the tendency to locate SO₂ monitoring stations in the vicinity of sources in order to ascertain compliance with air quality standards. The consequences of such considerations are shown in the time series for Site B (Figure 5b), which show observed SO₂ mixing ratios from two monitoring sites, one at the New Haven-Meriden, Connecticut site, 25 km from Site B, and one at the Waterbury, Connecticut site, 50 km from Site B (and 30 km from the Meriden site). The SO₂ mixing ratio at the New Haven-Meriden site exhibited rather large excursions and rather high values compared to the Waterbury site, which appears to be more regionally representative and at which mixing ratios were much suitable for comparison with the model calculations. A similar situation obtained for Site C, coastal Maine, for which the proximate sites exhibited highly varying temporal profiles of SO₂ (Figure 5c); for example, the Oxford, Maine, site, ~225 km from Site C, showed magnitudes closer to those obtained in the present calculations than did the more urban Portland, Maine, site ~175 km from Site C (and 100 km from the Oxford site). For Site A the only proximate stations, one in nearby Sioux Falls, South Dakota, 40 km from Site A, and the other in Mason City, Iowa, \sim 250 km from Site A, again exhibited rather differing SO₂ profiles (not shown). In summary the limited available comparisons suggest that the SO2 mixing ratios obtained from the model and used in calculation of H₂SO₄ production rate are regionally representative.

[54] Examination of vertical profiles of NPF-related variables yields insight into the dependence of key processes on altitude and the role of vertical mixing. Vertical profiles of [H₂SO₄], [NH₃], T, RH, $J_{\rm NPF}$, $N_{\rm ATK}$ and $D_{\rm g,ATK}$ at local noon averaged for July at Site B (which is at sea level) are shown for the different model variants in Figure 6. Sulfuric acid was formed at substantial concentrations up to 5 km, whereas ammonia was primarily below 2 km. Air temperature decreased monotonically to less than -40°C near the tropopause (near 12 km), whereas RH peaked near the top of the planetary boundary layer (PBL) at $\sim 1-1.5$ km. Under these conditions, appreciable NPF occurred at heights up to 5 km for EM97 and NIIR, whereas it was confined to the PBL for THN. NPF from BHNJ occurred at heights of between 1 and 3 km following the behavior of RH, and near the tropopause, whereas NPF from BHNV was negligible over the heights shown. $N_{\rm ATK}$ and $D_{\rm g,ATK}$ tended to be rather uniform throughout the PBL for several of the formulations, including BHNJ, for which there was

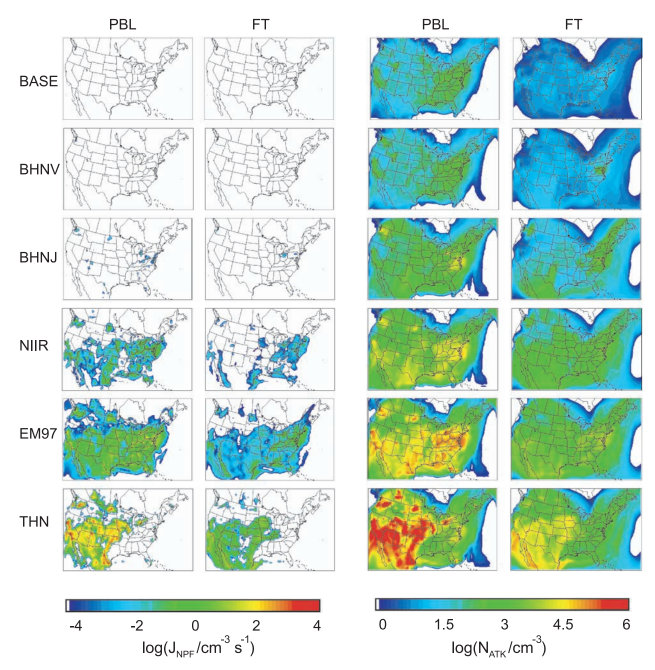


Figure 8. Spatial distribution of daily average J_{NPF} and N_{ATK} in the planetary boundary layer (PBL) and in the free troposphere (FT) for the different model variants on 20 July 2004.

little production at the surface, implying that substantial vertical mixing occurred within the PBL.

[55] The influence of the several NPF formulations on $J_{\rm NPF}$ and $N_{\rm ATK}$ is examined as a function of location over the entire continental United States in Figure 8, which shows the averages of these quantities over the PBL and over the free troposphere (FT). The dissimilar spatial distribution of $J_{\rm NPF}$ for different NPF formulations is attributed mainly to the spatial distribution of NPF precursors, NH₃ and H₂SO₄. Despite ammonia emissions being greatest over the middle and eastern United States (Figure 3), [NH₃] was lower in the eastern United States than in the western United States because of neutralization by sulfuric acid. In the western United States, because of lower emission of SO₂ (Figure 3),

much ammonia remained in the gas phase, allowing NPF by THN. As production of H₂SO₄ required time for the photochemical reaction, NPF by the EM97 and NIIR formulations were somewhat displaced from sources of SO₂, which were mostly located over eastern United States. Because SO₂ has a longer residence time than NH₃, EM97 and NIIR resulted in more widespread distributions of NPF than did THN, but at a lower rate. Widespread NPF events have been observed at a variety of places in the United States (for comprehensive reviews, see *Kulmala et al.* [2004] and *Yu et al.* [2008]), including locations that are rural (Idaho Hill, Colorado [*Marti et al.*, 1997]), coastal (Washington state coast [*Hegg et al.*, 1992]), and urban (e.g., Atlanta, Georgia [*Woo et al.*, 2001; *Stolzenburg et al.*, 2005], and Pittsburgh [*Stanier et al.*,

2004]). Although the fundamental mechanism explaining the ubiquitous background of ultrafine aerosols in the atmosphere remains poorly understood, examination of Figure 8 suggests that it might be better accounted for by superposition of several NPF mechanisms rather than by a single mechanism.

3.2. Sensitivity to Primary Emissions

[56] The sensitivities of modeled number concentrations and other aerosol properties to sizes (numbers) of emitted particles were examined by analyzing model results for the lowest level at Site B for three cases. In the D_0 case, the geometric mean diameters of emitted particle mass for the Aitken and accumulation modes ($D_{g,ATK}$ and $D_{g,ACC}$, respectively) retain their initial values of 30 nm and 300 nm, respectively. In the D/2 case, $D_{g,ATK}$ and $D_{g,ACC}$ were decreased to 15 nm and 150 nm, respectively, and in the $D\times 2$ case, $D_{g,ATK}$ and $D_{g,ACC}$ were increased from their initial values to 60 nm and 600 nm, respectively, while in both of the latter two cases the total emitted mass was held constant and distributed as noted above, with 99.9% going into the accumulation mode and 0.1% into the Aitken mode. Here examination was restricted to NPF occurring jointly by both the THN and the NIIR formulations (THN+NIIR). Time series of $[H_2SO_4]$, J_{NPF} , number concentrations in the Aitken and accumulation modes, and other related quantities for each case at the lowest model level at Site B are shown in Figure 9.

[57] A decrease (increase) in particle diameter by a factor of 2, for constant mass emissions, results in an increase (decrease) in the number of emitted particles by a factor of 8, and an increase (decrease) in the total concentration of surface area of emitted particles by a factor of 2; lines denoting these factors are shown in Figure 9. Associated with the increase (decrease) in surface area concentration would be a nearly proportional increase (decrease) in rate of condensation were no other factors at play. The situation with coagulation is more involved, as the rate of coagulation of particles of two diameters is directly proportional to the product of their number concentrations, the sum of their diameters, and the sum of their diffusion coefficients (each of which is inversely proportional to the square of the particle diameter for particles with diameters appreciably less than the mean free path of air, \sim 65 nm at sea level, and inversely proportional to the particle diameter for appreciably larger particles). If no other processes acted, a decrease (increase) in particle diameter by a factor of 2, for constant mass emissions, would result in an increase (decrease) in the first factor by $64 (= 8^2)$, in the next by 1/2, and in the third by 4 for self-coagulation of Aitken mode particles, resulting in an increase (decrease) of 128. The increase (decrease) for the self-coagulation rate of accumulation mode particles would be somewhat less because of their larger size and weaker dependence of diffusion coefficient on particle size. Lines denoting a factor of 128 in the self-coagulation rate for the Aitken and accumulation modes are also shown in Figure 9.

[58] Not surprisingly, the D/2 case systematically resulted in an increase, and the $D\times 2$ case in a decrease, in $N_{\rm ACC}$ (Figure 9), indicating that this quantity is largely determined by primary emissions. However, the increase (decrease) in $N_{\rm ACC}$ was typically about a factor of 4, and the decrease

(increase) in $D_{g,ACC}$ was slightly less than a factor of 2, less than what would have resulted if the number concentration were controlled entirely by primary emission and no coagulation occurred (a factor of 8 for N_{ACC} and a factor of 2 for $D_{g,ACC}$). The quantities [SO₄]_{ACC}, [NH₄]_{ACC}, and [SOA]_{ACC} were nearly the same for each case at almost all times, although in some instances (e.g., 21 July) $[SO_4^{2-}]_{ACC}$ and $[NH_4^+]_{ACC}$ were slightly greater for the D/2 case and slightly less for the $D\times 2$ case than for the D_0 case, reflecting the increased (decreased) rate of condensation associated with the increased (decreased) surface area concentration accompanying the decrease (increase) in size of emitted particles. Additionally, there was a nearly uniform increase (decrease) in the self-coagulation rate of accumulation mode particles by roughly a factor of 50 accompanying the decrease (increase) in the diameter of emitted particles. As expected, the rate of wet deposition of accumulation mode particles was greatest for the D/2 case, reflecting the greater number of particles available to be removed by precipitation.

[59] In contrast to these results, N_{ATK} was generally less for the D/2 case, and slightly greater for the $D\times 2$ case, than for the D_0 case (Figure 9), implying that primary emissions were not a dominant factor in controlling N_{ATK} . Additionally, $D_{g,ATK}$ was often less for the D/2 case than for the D_0 case, but it was nearly the same for the $D\times 2$ and D_0 cases. These results can be explained by suppression of NPF by the greater condensational sink (in both modes) for H₂SO₄ accompanying the decrease in size of emitted particles (i.e., D/2 case), resulting in a decrease in [H₂SO₄], this increased condensation sink and consequent decrease in [H₂SO₄] providing less favorable conditions for NPF. For example, $D_{g,ATK}$ was the greatest in the $D\times 2$ case on 12 July (green circle in Figure 9) when $J_{\rm NPF} < 10^{-3} \, {\rm cm}^{-3} \, {\rm s}^{-1}$, whereas $D_{\rm g,ATK}$ was the smallest in the same case on 15 July (brown circle in Figure 9) when $J_{\text{NPF}} > 10^2 \text{ cm}^{-3} \text{ s}^{-1}$. In support of this explanation, there were sometimes large differences in J_{NPF} between the several cases with less NPF typically occurring for the D/2 case than for the other two cases. Also, $F_{\rm KK}$ was often greater for the $D\times 2$ case than for the others, and there were several instances in which IMT occurred for the $D\times 2$ case but not either of the others. These findings support the conclusion that $N_{\rm ATK}$ is determined much more by NPF than by primary emissions. Also consistent with this explanation is the typically greater rate of self-coagulation in the Aitken mode in the $D\times 2$ case than in the D/2 case. $[SO_4]_{ATK}$ and $[NH_4]_{ATK}$ were occasionally greater for $D\times 2$ case, although they were often independent of case. The dry deposition flux of Aitken mode particle number, for which the strong peaks tend to accompany large values of N_{ATK} , was also nearly independent of case.

3.3. Budgets of Aerosol Number Concentration

[60] The factors controlling aerosol number concentrations in the Aitken and accumulation modes are further examined by consideration of the number budget, that is, the magnitudes of the various sources and sink rates of aerosol number concentration, in both modes, averaged over the entire modeling domain and time for each model variant (Figure 10). The overwhelmingly dominant source of number concentration in the Aitken mode was NPF, which greatly exceeded primary emissions in all model variants (except the

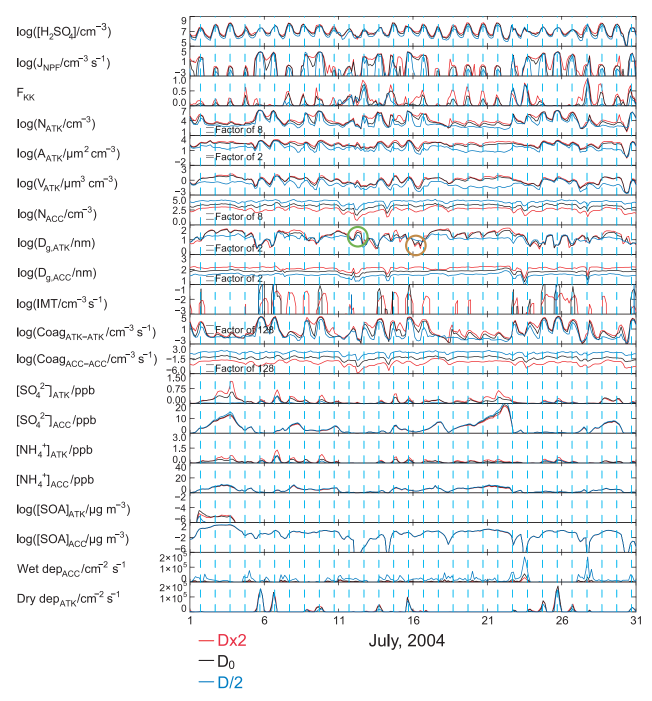


Figure 9. Time series of [H₂SO₄], $J_{\rm NPF}$, $F_{\rm KK}$, $N_{\rm ATK}$, $A_{\rm ATK}$, $V_{\rm ATK}$, $N_{\rm ACC}$, $D_{\rm g,ATK}$, $D_{\rm g,ACC}$, rate of IMT of aerosol number, intramodal coagulation rates for the Aitken and the accumulation modes, dry-air molar mixing ratios of sulfate and ammonium in the Aitken and accumulation modes, mass concentrations of SOA in the Aitken and the accumulation modes, wet deposition flux of accumulation mode particle number, and dry deposition flux of Aitken mode particle number at the lowest level (0–20 m above the ground) for THN + NIIR at Site B during July 2004, with mass emissions at $D_{\rm g,ATK}$ = 30 nm and $D_{\rm g,ACC}$ = 300 nm (D_0 case), and for $D\times 2$ and D/2 cases. The quantities $D_{\rm g,ATK}$, $D_{\rm g,ACC}$, $A_{\rm ATK}$, and $V_{\rm ATK}$ are reported for dry particles, although for use in evaluation of coagulation rates, etc., they were calculated for particles at their ambient (i.e., wet) sizes. Abundances of [SO₄²⁻] and [NH₄⁺] are presented as dry-air molar mixing ratios in nmol mol⁻¹ (ppb, parts per billion); at 298 K and 1 atm, a molar mixing ratio of 1 ppb is equivalent to a mass concentration of 3.9 μ g m⁻³ for sulfate and to 0.74 μ g m⁻³ for ammonium. Lines denoting factor of 8 in $N_{\rm ACC}$ and $N_{\rm ATK}$, factor of 2 in $D_{\rm g,ACC}$, $D_{\rm g,ATK}$, and $A_{\rm ATK}$, and factor of 128 in intramodal coagulation rates Coag_{ATK-ATK} and Coag_{ACC-ACC} are shown. Green circle represents instance of $D_{\rm g,ATK}$ being largest in the $D\times 2$ case; brown circle represents instance of $D_{\rm g,ATK}$ being largest in the $D\times 2$ case;

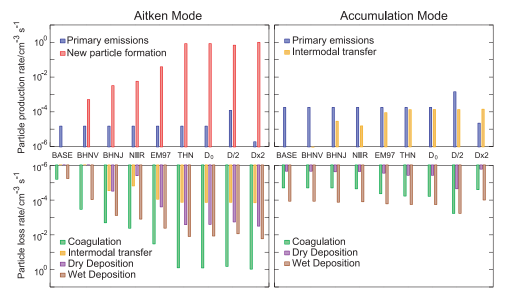


Figure 10. Production and loss rates of aerosol particle number concentration in the Aitken and accumulation modes by different processes for the different model variants averaged over the entire modeling domain and time period.

BASE case, for which there was no NPF); consequently, N_{ATK} is insensitive to the specific fraction of mass assumed to be emitted into the Aitken mode, here taken as 0.1%. The main sink of number concentration in the Aitken mode was coagulation, with minor contributions from wet and dry deposition. IMT was generally negligible, being no more than a few percent of coagulation for BHNJ, and even less for other model variants. Although J_{NPF} differed by more than 3 orders of magnitude among the several NPF formulations (Figure 10), the resulting N_{ATK} exhibited a range of only about 2 orders of magnitude (Table 2), the smaller range being attributed to increased coagulation of the very small particles resulting from larger values of J_{NPF} . The turnover time for particle number in the Aitken mode, $\tau_{\mathrm{N,ATK}}$, defined as the total number of Aitken mode particles in the domain divided by the total loss rate due to physical processes (i.e., not including transport in and out of the domain), ranged from 0.03 d to 0.5 d among the five NPF formulations (Table 2), being determined largely by the coagulation rate. The smallest turnover times occurred for THN and THN + NIIR (i.e., D_0 , $D \times 2$, and D/2), as these resulted in the largest values of $N_{\rm ATK}$, and thus even greater coagulation rates, which are directly proportional to the squares of the number concentrations.

[61] The dominant source of aerosol number concentration in the accumulation mode was primary emission, although for some model variants IMT provided a nearly equal contribution (Figure 10). Thus, the largest variation of $N_{\rm ACC}$ occurred by altering the size of emitted particles (Table 2). As IMT introduced particles into the accumulation mode at small sizes, the overwhelming contribution to the mass concentration in this mode was provided by emissions. The dominant sink of number concentration in this mode was wet deposition (including removal of unactivated particles by coagulation on activated accumulation mode particles), which was nearly the same for all model variants except the D/2 case, with coagulation generally providing a relatively minor contribution, which was also roughly the same for all model variants except the D/2 case. For the D/2 case both contributions were nearly

equal to each other and greater than for the other model variants. In this case the initial particles were smaller and IMT resulted in more and smaller particles in this mode. The contribution from dry deposition was generally negligible for all model variants. Consequently, the turnover time for particle number in the accumulation mode, $\tau_{\rm N,ACC}$, was nearly independent of model variant at around 2.7 d, except for the D/2 case when it was 1.6 d because of the increased loss of particle number due to coagulation (Table 2).

[62] The mean turnover time for aerosol mass, which resides almost entirely in the accumulation mode (Figure 5), is defined similarly to that for aerosol number, although loss mechanisms differ. Importantly, coagulation results in a loss of number but leaves mass unchanged; thus, only wet and dry deposition act as sinks for aerosol mass. However, the importance of these processes differs between mass and number concentrations, as they affect particles of different sizes differently. The turnover time for total aerosol mass was ~ 8 d for all model variants in the present study, characteristic of the time between precipitation events (Figure 5). This time is greater than that for aerosol number, for which coagulation is also a sink, and is comparable to

Table 2. Time- and Domain-Averaged Number Concentrations in the Aitken and Accumulation Modes and Their Turnover Times for All Model Variants

	Aitken Mode		Accumulation Mode	
Model Variant	$N_{\rm ATK}/{\rm cm}^{-3}$	$ au_{ m N,ATK}/{ m d}$	$N_{\rm ACC}/{\rm cm}^{-3}$	$ au_{ m N,ACC}$ /d
BASE	3	2.8	33	2.7
BHNV	20	0.51	34	2.7
BHNJ	139	0.54	39	2.7
NIIR	99	0.20	36	2.7
EM97	517	0.15	48	2.6
THN	2581	0.035	60	2.8
D_0	2436	0.033	60	2.7
D/2	1863	0.031	169	1.6
$D \times 2$	3251	0.037	35	3.1

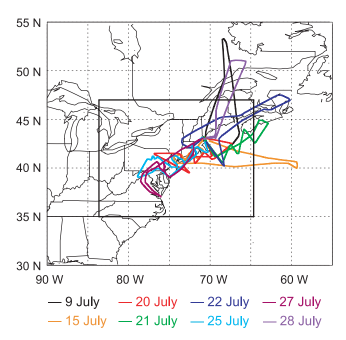


Figure 11. Flight paths of the NOAA WP-3D aircraft during the International Consortium for Atmospheric Research on Transport and Transformation (ICARTT) 2004 study. Inner box denotes the domain of the 12-km resolution model.

that of sulfate mass reported in several other model studies (e.g., 6.5 d by *Adams and Seinfeld* [2002] and 7.0 d by *Benkovitz et al.* [2004]), although it is considerably greater than the mean of 4.1 ± 0.7 d for 15 studies reviewed by *Textor et al.* [2006]. The longer turnover time found here compared to those studies is attributed to the paucity of precipitation events in the model domain during July 2004.

4. Comparison With Observations

[63] Model results were compared with aircraft measurements taken over the northeast United States and eastern Canada using the National Oceanographic and Atmospheric Administration (NOAA) WP-3D (P3) aircraft during the ICARTT 2004 study [Fehsenfeld et al., 2006; Nowak et al., 2007]. Tracks of the flights used in the present comparisons are shown in Figure 11 and are seen to include Sites B, C, and D. For each observational data point shown in Figure 12, data from the eight nearest grid-cell midpoints to the location of the P3 were sampled and then bilinearly interpolated to the location of the aircraft. Model calculations of number, surface area, and volume concentrations for the Aitken and accumulation modes for the several NPF formulations were summed and compared to the integrals of the aircraft measurements for (dry) diameters from 3 nm to 2.5 μ m. Although the number concentration was dominated by the Aitken mode, the main contributions to the concentrations of surface area and volume were from the accumulation mode. The different NPF formulations differed primarily in number concentrations, and exhibited nearly the same concentrations of surface area and volume, similar to the situation with the model results at the different sites at ground level (Figure 5).

[64] In some instances the model did poorly (some of these are shown as orange bands in Figure 12), especially at higher altitude and during the ascent and descent of the aircraft. There seem to be several reasons for the poor performance. For example, on 9 July the aircraft encountered a biomassburning plume at 4 km, which is thought to have come from Canadian and Alaskan forest fires [Fehsenfeld et al., 2006]; the wind in both observation and model was from the north where no Canadian area sources were taken into account. The sharp peaks in surface area and volume concentrations observed at height 3 km from 1700 to 1800 UST on 20 July came from a biomass-burning plume [Fehsenfeld et al., 2006], which was not captured by the model. A biomassburning plume was also encountered on 28 July [Fehsenfeld et al., 2006]. High surface area and volume concentrations in the lower marine atmosphere (61°W, 40.5°N) between 1500 and 1600 UTC on 15 July may have been due to the presence of sea salt particles, emission of which was not treated in the model. The modeled wind speed was relatively high (\sim 8 m s⁻¹ at 7 m) because the area was under the influence of a cyclonic low-pressure system, and production of sea salt particles would thus be expected. There were probably other factors that also contributed, as at times immediately following these the concentrations of surface area and volume simulated by the model agreed fairly well with observations although the number concentrations were underestimated by nearly an order of magnitude.

[65] In general, however, the model did rather well in reproducing aerosol concentrations, especially at lower altitudes (some instances are denoted by brown circles in Figure 12). For example, the rather fresh Boston plume on 9 July around 1800 UTC was captured by the model, as was a well defined New York City plume on 15 July from 1800 to 2100 UTC [Fehsenfeld et al., 2006]. In addition, the high values of the number, surface area, and volume concentrations observed on 20 July, reaching up to 2 km, were well matched by the modeled ones through all heights. The model also performed well on 25 and 27 July. Among the several NPF formulations considered EM97 reproduced number concentration best, but agreement was good only at lower altitudes.

[66] A substantial fraction of the differences between model and observation may be due to subgrid variation [Benkovitz et al., 2004]. The effect of subgrid variation was examined by comparing results from the two different model resolutions, 60 km with 3 h output and 12 km with 10 min output, both using the sum of the THN and NIIR formulations, with P3 measurements taken on 20 July (Figure 13). For most of the quantities compared, the results from the two resolutions were quite similar, and for some quantities they agreed rather well with the observations. Modeled values of $[SO_4^{2-}]$ were greater than observations, but the 12-km resolution version of the model generally provided greater accuracy, as it did also for [NO₃]. The high temporal variability of modeled [NO₃] between 1430 and 1500 UTC in the 60-km resolution version of the model may be due to sampling model data from the edge of the modeled plume centered at 72°W and 39°N (not shown here). Both resolutions underestimated [NH₃], but as [NH₃] is, to good approximation, equal to the difference (in equivalent units) between total [N(-III)] (the sum of $[NH_3]$ and $[NH_4^+]$) and the sum of $[SO_4^{2-}]$ and $[NO_3^-]$,

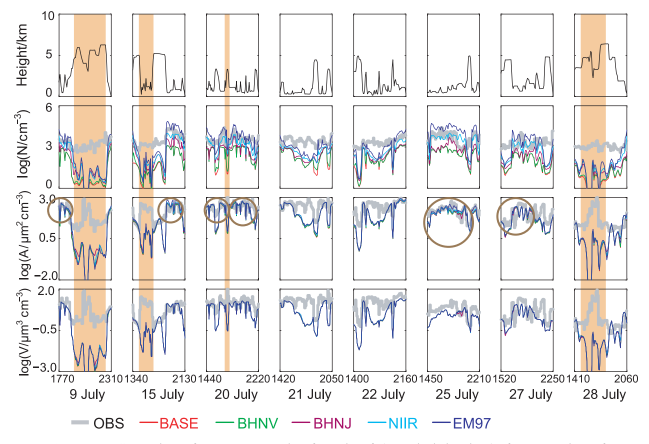


Figure 12. Comparison of measurements taken from aircraft (gray shaded regions) of concentrations of number N, surface area A, and volume V with results of different model variants for eight selected days during the ICARTT 2004 study. Measured concentrations are for (dry) diameters from 3 nm to 2.5 μ m; modeled values are calculated using ambient (wet) diameters but reported for dry diameters, as the sum of the concentrations in the Aitken and accumulation modes. Height refers to height above mean sea level. Orange bands denote some period of poor performance, and brown circles denote some periods of good performance.

underestimation of [NH₃] is mainly a consequence of the overestimation of sulfate, which occurs in both model resolutions (although considerably less so in the higher resolution model). The low-biased [NH₃] may be the reason that the model underpredicted aerosol number concentration, exhibiting much lower concentrations of small particles (Figure 13, last three panels) between 1630 and 1700 UTC and between 1800 and 1930 UTC than those measured, which attain values up to near 2×10^4 cm⁻³. Concentrations of number, surface area, and volume were often underestimated by both model resolutions, although in some instances (e.g., 1930–2000 UTC) the 12-km model overpredicted number concentration but underpredicted surface area and volume concentrations. One interesting feature of the comparisons is the relatively good agreement between measured and modeled SOA, implying that it might not be underestimated in models as suggested by some investigators.

5. Summary and Conclusions

[67] The dependence of aerosol number concentration and other intensive and extensive aerosol properties on different formulations of new particle formation NPF and

on the particle size and number of primary emissions has been examined using the regional CMAQ model over the continental United States for July 2004. The calculated rate of NPF, $J_{\rm NPF}$, varied greatly for the several NPF formulations examined, and for any given NPF formulation J_{NPF} varied greatly in time and space. Because of the relatively high ammonia concentration in the western United States compared to the eastern United States, where high SO2 emissions and resultant high sulfate concentrations result in low NH₃ concentrations, ternary homogeneous nucleation (THN) involving ammonia together with sulfuric acid and water was the dominant NPF mechanism in the western United States. In the eastern United States, J_{NPF} was greatest for the empirical formulation of Eisele and McMurry [1997]. New particle formation by ion-ion recombination (NIIR) provided a substantial contribution across the whole domain; however, recent field studies showing that ioninduced nucleation is insignificant compared to other mechanisms of NPF [Eisele et al., 2006; Iida et al., 2006] suggest that the NIIR formulation employed here may substantially overestimate $J_{\rm NPF}$ by that mechanism. NPF by THN and by NIIR occurred primarily in the planetary boundary layer, whereas for both binary (sulfuric acid and water) homoge-

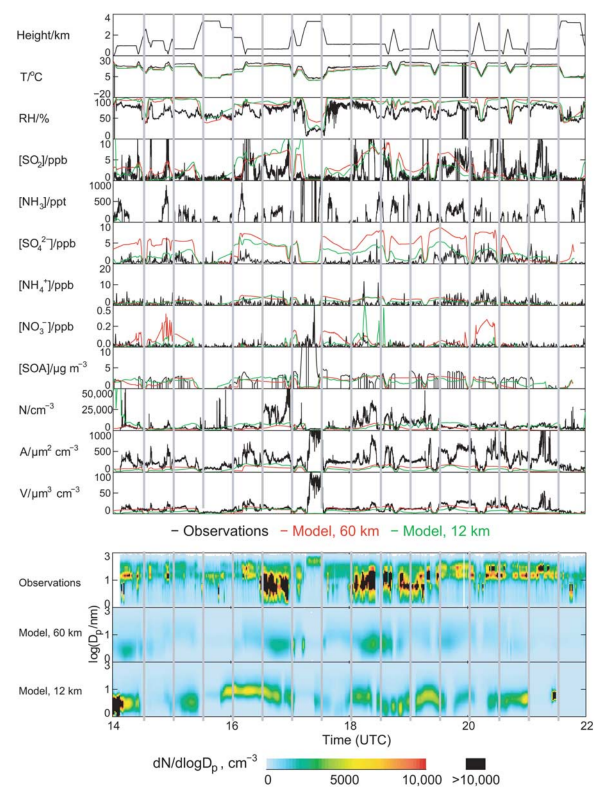


Figure 13. Comparison between measurements taken from aircraft and model simulations for both the 60-km resolution and the 12-km resolution for THN + NIIR on 20 July 2004. Concentrations of gaseous species are given as dry-air molar mixing ratios in nmol mol⁻¹ (ppb, parts per billion), except for [NH₃], which is in pmol mol⁻¹ (ppt, parts per trillion). Measured concentrations of particle number N, surface area A, and volume V are for (dry) diameters from 3 nm to 2.5 μ m; modeled values are calculated using ambient (wet) diameters but reported for dry diameters, as the sum of the concentrations in the Aitken and accumulation modes. Height refers to height above mean sea level. The last three panels compare the size distribution of aerosol number concentration for diameter from 2 nm to 1 μ m.

nous nucleation formulations considered NPF occurred primarily near the tropopause because of the low temperature there, but at a relatively low rate.

[68] For each NPF formulation considered the major source of aerosol number concentration in the Aitken mode was NPF despite the large variation in J_{NPF} among the several formulations, with the contribution from emissions being considerably less. The major source of domainaverage aerosol number concentration in the accumulation mode was primary emission, but for instances of intense NPF a comparable source was provided by transfer of Aitken mode particles to the accumulation mode, although this process contributed very little to the accumulation mode mass concentration. When NPF occurred, the characteristic diameter of Aitken mode particles $D_{g,ATK}$ exhibited a diurnal variation opposite to that of J_{NPF} because NPF reduced $D_{g,ATK}$, whereas subsequent coagulation increased $D_{\rm g,ATK}$ to near its previous value. In contrast, the characteristic diameter of accumulation mode particles $D_{\rm g,ACC}$ exhibited little dependence on NPF formulation, being controlled mainly by the diameter of emitted particles.

[69] The sensitivity of aerosol properties to change in size of emitted particles for constant mass emission rate was examined, a factor of 2 decrease (increase) in particle size resulting in a factor of 8 increase (decrease) in number emission rate. Surprisingly, it was found that an increase (decrease) in the number of emitted particles resulted in a decrease (increase) in the number concentration of Aitken mode particles N_{ATK} . This is attributed to NPF being controlled by sulfuric acid vapor; when the size of emitted particles is decreased, the resultant increase in surface area increases the sink rate of sulfuric acid vapor, suppressing NPF and resulting in an overall decrease in N_{ATK} . In contrast, the dominant sensitivity of the number concentration of accumulation mode particles N_{ACC} is to the size (number) of the emitted particles (for constant mass emissions). The change in N_{ACC} in response to changes in number and size of emitted particles was in the direction expected, but at a somewhat lower magnitude (roughly a factor of 4) than would be expected (a factor of 8) if primary emitted particles were the only source of accumulation mode particles and no coagulation occurred.

[70] The sensitivity of N_{ACC} to NPF formulation and to the size (number) of emitted primary particles would result in a high uncertainty in the calculation of aerosol indirect effects, which depend on aerosol number concentration. In this context, it might be noted that an increase of only 30% in droplet number concentration in marine stratus clouds globally corresponds to a climate forcing of 1 W m⁻² [Charlson et al., 1992]. The present study thus demonstrates the need for both NPF and primary emissions to be better constrained if the direct and first indirect effects are to be accurately represented in models of aerosol forcing and ultimately in climate models.

[71] Comparison of model results to measurements yielded many instances of rather good performance of the model for surface area and volume, especially in the planetary boundary layer, where emissions were mainly within the model domain. However, in the free troposphere the model accuracy was often less; this is attributed to missing sources of particles and/or long-range transport from outside the model domain. This finding suggests the

need for addition of biomass burning sources and for more realistic boundary conditions for regional-scale models, or perhaps extension of the model domain to a larger scale; the need to include the entire northern hemisphere in modeling sulfate aerosol mass concentration has been noted previously [Benkovitz et al., 2004]. Chemical composition and number concentrations obtained from the 12-km resolution version of the model were closer to measurements than were those obtained from the 60-km resolution version.

[72] The main limitations of this model, which might affect the analysis of the consequences of different treatments of NPF and emissions, are the modal framework (specifically the constrained aerosol evolution resulting from the assumption of lognormal size distributions and the artificial intermodal transfer from the Aitken mode to the accumulation mode), inaccuracy in the conversion factor from nuclei to new particles due to neglect of self-coagulation, uncertainty in the size of emitted primary particles, and lack of emissions of aerosol species such as sea salt and biomass burning. A potential concern with the modal method as applied here is the inclusion of newly formed particles in the Aitken mode and the potential resultant broadening of this mode. Although another mode containing newly formed particles might be added, this approach would necessitate deciding when sufficient new particle formulation occurred to introduce this mode, establishing criteria for intermodal transfer between this new particle mode and the Aitken mode, and the like. In any event, as the modeled geometric standard deviation of the Aitken mode σ_{ATK} exhibited values between \sim 1.4 and 2.1, rarely exceeding 2.0, such broadening would seem to be of minimal concern.

[73] While this study points to and quantifies some of the consequences that result from differing assumptions on NPF formulation and on primary emissions, model studies such as this can at best determine only the consequences of the assumptions that are coded in the model. Resolution of the large differences in number concentration revealed here for different model variants ultimately rests on measurements. In that respect we note the dearth of measurements characterizing size distributions (and even more so, size-distributed composition) of primary particulate matter from the many processes that give rise to such emissions. Characterization of particulate emissions has focused almost entirely on mass emission rates, this focus arising from air quality considerations and standards that are prescribed in terms of mass concentrations of particulate matter with (aerodynamic) diameter (at a given RH) less than a given value, for example, 2.5 μ m or 10 μ m in the United States. Consequently assumptions such as that made here that 99.9% of the emitted mass is in the accumulation mode and only 0.1% is in the Aitken mode, have little support in measurement (although the results of this investigation suggest that the specific fraction of mass assumed to be emitted into the Aitken mode has little effect on N_{ATK}). Likewise, NPF rates and their dependence on controlling variables are highly uncertain, and even the mechanisms involved are not completely understood. Here we treated only NPF involving sulfuric acid, ammonia, and water vapor, which have been thought to be the major contributors to NPF in the atmosphere and consequently received considerable attention. However, evidence is becoming available from measurements that organic substances can also participate in NPF. For example, measurements of the composition of newly formed particles with diameter as low as 10 nm in Mexico City [Smith et al., 2008] have shown that these particles may contain an order of magnitude more organics than sulfates, with growth rate based on measured sulfuric acid concentration being an order of magnitude less than the observed growth rate. Additionally, parameterizations for NPF rates are being updated on the basis of new studies and as other mechanisms are considered. For instance, recent investigations [Kuang et al., 2008] suggest that the exponent p in equation (3) denoting the power law dependence of nucleation rate on sulfuric acid concentration is equal to 2 to within experimental uncertainty, and that the value of K used in equation (3) (denoting the magnitude of the rate) is typically greater than that used in the model by more than an order of magnitude. An updated formulation of ternary homogeneous nucleation of sulfuric acid, ammonia, and water [Merikanto et al., 2007] that includes presumed effects of stable ammonium bisulfate (despite absence of experimental evidence for the presence of this substance) yields nucleation rates several orders of magnitude lower than those of Napari et al. [2002]. A new formulation for the fraction of stable nuclei that grow to become new particles, the quantity F_{KK} in equation (2), was recently presented by Lehtinen et al. [2007]; however, there remains much uncertainty in this fraction, and the assumptions required for its computation, that the growth rate and condensation sink remain constant during a time step, and that self-coagulation of nuclei smaller than new particles does not occur, preclude the possibility of the model capturing an extremely strong pulse-type nucleation that is self-quenching. Confident representation of the responsible processes in models, and the dependence on controlling variables (concentrations of precursor gases, insolation, and the like) requires determination of these dependences in laboratory studies and confirmation of these mechanisms by field measurements.

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